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Modelling a Linker Mix-and-Match Approach for Controlling the Optical Excitation Gaps and Band Alignment of Zeolitic Imidazolate Frameworks

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Abstract: Tuning the electronic structure of metal–organic frameworks is the key to extending their functionality to the photocatalytic conversion of absorbed gases. Herein we discuss how the band edge positions in zeolitic imidazolate frameworks (ZIFs) can be tuned by mixing different imidazole-based linkers within the same structure. We present the band alignment for a number of known and hypothetical Zn-based ZIFs with respect to the vacuum level. Structures with a single type of linker exhibit relatively wide band gaps; however, by mixing linkers of a low-lying conduction edge with linkers of a high-lying valence edge, we can predict materials with ideal band positions for visible-light water splitting and CO₂ reduction photocatalysis. By introducing copper in the tetrahedral position of the mixed-linker ZIFs, it would be possible to increase both photo-absorption and the electron–hole recombination times.

The attractive pore structure of metal–organic frameworks (MOFs) has led to their huge success in gas adsorption and separation applications.^[1] Recently, there has been growing interest in extending the uses of MOFs to photocatalysis, as this would allow, for example, using a single material for both capture and conversion of CO₂ in a “one-pot” approach. The chemical diversity of MOFs can be exploited to tailor their electronic structure to the particular photocatalytic application, whereas their porous network can provide access of reactant molecules to the active sites.^[2] Some success has been achieved on MOF-based photocatalysis in the last few years, typically involving MOFs with metal nanoparticles or complexes incorporated inside their pores, or introduced via post-synthetic ligand modification.^[3] But there have been few reports where MOFs (without co-catalysts) have shown intrinsic photocatalytic activity. Gascon et al. reported the use of the IRMOF family of MOFs, with band gaps tuned via linker functionalization, in photocatalytic propylene epoxidation,^[4] while Gomes-Silva et al. have shown the potential of UiO-66 and related MOFs for photocatalytic hydrogen generation.^[5] MOFs with intrinsic photocatalytic behavior would indeed reduce the cost and complexity of catalyst preparation, but achieving the required electronic structure and catalytic behavior without the contribution from external species is challenging.

Herein we present the findings of our computational search for MOFs with the required electronic structure to operate as single-semiconductor, intrinsic photocatalysts in solar-fuel synthesis reactions. The search was focused on the zeolitic imidazolate frameworks (ZIF) family, as they exhibit the required high chemical and thermal stability in aqueous solution.^[6] ZIF structures with the same topology (SOD) and tetrahedral metal (Zn), but different imidazolate-based linkers or combination of linkers, were computationally generated. The SOD topology (Figure 1) was chosen because of its flexibility to accept a wide range of chemical compositions, in terms of both linkers and metals.^[6b] Recently, ZIFs with SOD topology have been experimentally investigated and shown promise in the context of photocatalytic applications.^[7] Also, its primitive cell is small enough to allow an efficient screening of a range of compositions using quantum-mechanical calculations.

We initially considered ZIF structures with composition ZnX₂ and 12 different linkers (X), including the bare imidazolate (Im) linker as well as the following functionalizations: added nitro (nIm), carboxaldehyde (Ica), or methyl

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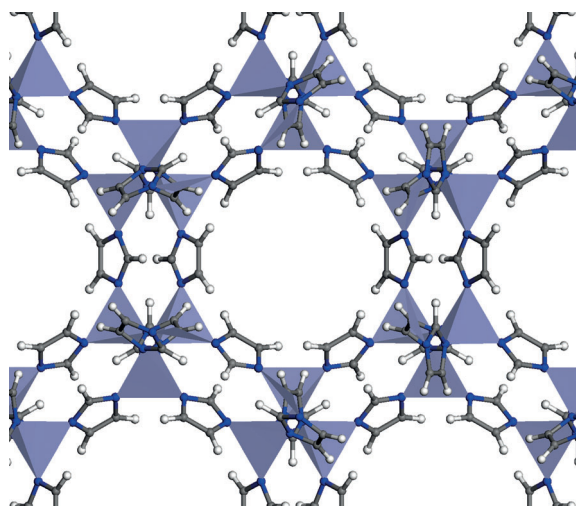


Figure 1. ZIF with composition ZnIm_2 and SOD topology, viewed along the direction of the largest pore. The tetrahedra are centered on Zn atoms; N blue, C gray, H white balls.

(mIm) side chains at ring position 2; di-cyanide (dcnIm) or di-chlorine (dclm) at positions 4,5; fused benzene (bIm), pyridine (abIm, acIm) or purine (pur) 6-membered rings; and fused thiophene (tIm) or furan (fIm) 5-membered rings (Figure 2a). ZIF structures with some of these linkers have been reported before, including Cu(Im)_2 , Zn(Ica)_2 , Zn(bIm)_2 , Zn(mIm)_2 and Co(nIm)_2 with SOD topology, as well as Zn(abIm)_2 and Zn(pur)_2 with LTA topology.^[6b] To date, no ZIFs have been synthesized with tIm and fIm linkers,

although these molecular units are reported in the patent literature.^[8]

The electronic structure was determined using density functional theory (DFT) with a screened hybrid density functional,^[9] and the electronic levels were aligned with the vacuum scale.^[10] The band edges in these molecular solids, which are better referred to as the highest occupied crystal orbital (HOCO) and lowest unoccupied crystal orbital (LUCO) levels, are shown in Figure 2b. The widest HOCO–LUCO gap is obtained in the case of the bare imidazole linker, that is, for Zn(Im)_2 (5.2 eV), while the narrowest gap corresponds to Zn(nIm)_2 (3.3 eV). The wide range observed illustrates the tunability of the electronic properties of ZIFs upon linker modification, which is consistent with previous theoretical and experimental findings for other MOFs families.^[11]

For applications in photocatalysis, not only the magnitude of the band gap but also the absolute positions of the band edges are important. For water splitting with a single-semiconductor photocatalyst, the band edges should straddle the redox potentials for water photolysis,^[12] that is, the HOCO should be below the energy level of the oxygen evolution reaction (OER: $\text{H}_2\text{O} \leftrightarrow 2\text{H}^+_{(\text{aq})} + \frac{1}{2}\text{O}_{2(\text{g})} + 2\text{e}^-$), and the LUCO should be above the energy corresponding to the hydrogen evolution reaction (HER: $2\text{H}^+_{(\text{aq})} + 2\text{e}^- \leftrightarrow \text{H}_{2(\text{g})}$). In the vacuum scale, and at pH 0, the HER level is located at -4.44 eV, and the OER level is located at -5.67 eV.^[13] At temperature T and pH > 0 , these levels are shifted up by $\text{pH} \times (k_{\text{B}}T \times \ln 10)$, where k_{B} is Boltzmann's constant. In the case of CO_2 conversion to fuels (e.g. methanol), the LUCO must be above the redox potential for the CO_2 reduction half-

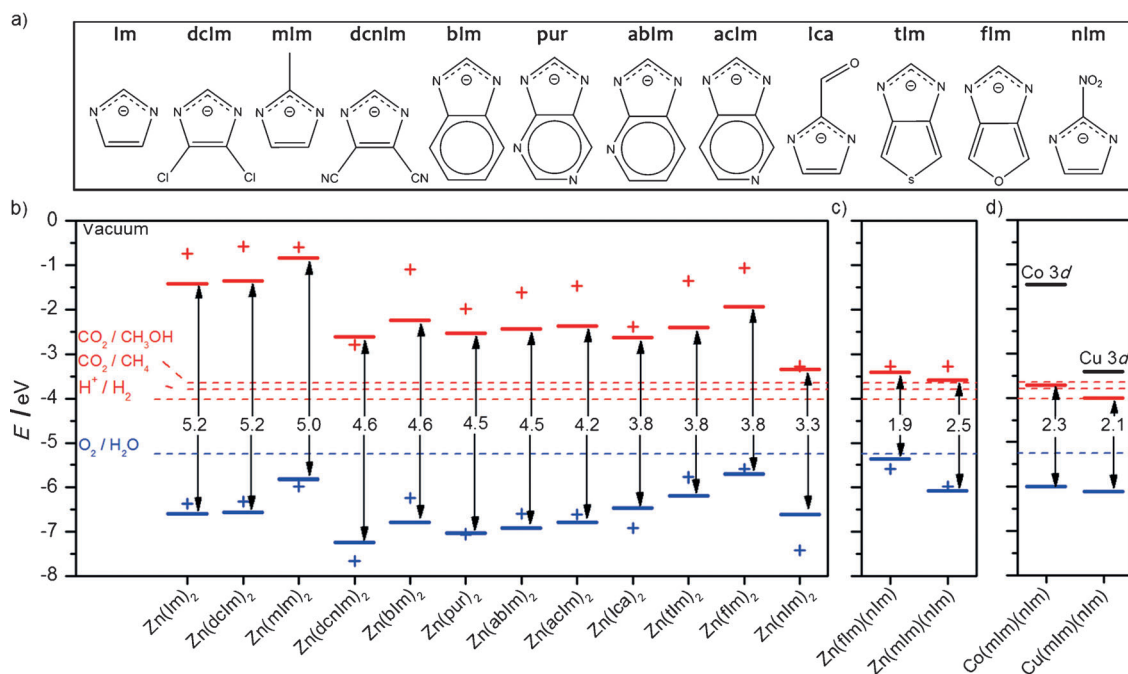


Figure 2. a) List of linkers (X) in the ZIF structures with composition ZnX_2 . b) Positions of the highest occupied energy levels (blue) and lowest unoccupied energy levels (red) of the ZnX_2 crystal structures (lines), and of the isolated HX molecules (crosses). The potentials for H_2O splitting and two different CO_2 reduction reactions are shown as dashed lines. c) Equivalent plot for the mixed ZIF structures created by combining nIm with mIm or fIm linkers; the crosses represent the higher HOMO (blue) and the lower LUMO (red) among the two mixed linkers. d) Equivalent plot for mixed mIm/nIm ZIF structures with Co or Cu centers.

reaction.^[14] Figure 2b shows that the HOCO and LUCO levels of the ZnX_2 ZIF structures considered here straddle the redox levels for both water splitting and methane/methanol synthesis. However, for these photocatalysts to work efficiently under solar radiation, a narrower band gap of around 2 eV is desirable, as this would allow the absorption of the visible range solar spectrum, which carries most of the solar radiation energy. Furthermore, the band gaps of these single-linker ZIFs correspond to intra-linker excitations, which is not convenient for photocatalysis, because some degree of electron-hole separation is needed to prevent fast recombination of the charge carriers.^[15]

We have therefore gone a step further and generated new hypothetical ZIFs by mixing different linkers within the same structure, to tailor the band edges to the required positions and to attain electron-hole separation via inter-linker excitations. We have chosen to mix a low-LUCO linker ($\text{X} = \text{nIm}$) with high-HOCO linkers ($\text{Y} = \text{Im}$, mIm , or flm) to form ZnXY structures with potential low-energy inter-linker excitations. In these 50:50 mixed-linker structures, two well-defined ordered configurations can be formed, considering the symmetry of the lattice. For each composition, the lowest-energy mixed configuration is significantly stable with respect to the separated pure phases (by 0.2–0.4 eV per formula unit, see Supporting Information), which indeed suggests they should be able to form experimentally. It is known that mixed-linker ZIFs can be synthesized solvothermally, via the reaction of equimolar amounts of the two protonated linkers with the divalent metal nitrate; this has been demonstrated by Yaghi et al.,^[16] who reported the formation of structures with ordered distributions of linkers.^[17] Figure 2c shows the HOCO and LUCO levels for the most stable mixed ZIFs with compositions $\text{Zn}(\text{flm})(\text{nIm})$ and $\text{Zn}(\text{mIm})(\text{nIm})$. They perfectly straddle the targeted redox potentials and lead to gaps of 1.9 and 2.5 eV, respectively, which are very convenient for applications involving solar-light absorption. The projected electronic density of states (DOS) in the mixed ZIFs (see Supporting Information) confirms that the LUCO is mainly contributed by the nitro group in the nIm linker, while the HOCO is contributed by the second linker.

This linker “mix-and-match” method can then be used more generally to target desired electronic structures of ZIFs for photocatalytic applications. The reason why the procedure works becomes apparent from examining the energies of the highest-occupied (HOMO) and lowest-unoccupied (LUMO) molecular orbitals of the isolated linkers. The HOMO and LUMO of the neutral molecules HX are represented as crosses in Figure 2b (and also in Figure 2c, but using the higher HOMO and the lower LUMO among the two mixed-linkers). Despite some fluctuations, the HOCO/LUCO levels in the ZIFs follow the same trend as the HOMO/LUMO levels in the corresponding gas-phase linker molecules. As Figure 3 shows, there is an excellent correlation between the excitation gaps in the linkers and those in the corresponding ZIFs. The correlation extends to the mixed-linker ZIFs, if we take the difference between the higher HOMO and the lower LUMO between the two linkers. It is clear from this analysis that the positions of the band edges in the crystalline material are determined, to a large extent, by

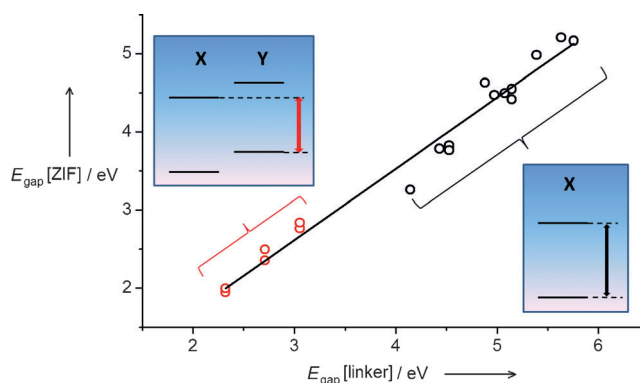


Figure 3. Correlation between the HOCO–LUCO gaps of the ZIF solids (ZnX_2 or ZnXY) and the HOMO–LUMO gaps of the protonated linkers in gas phase. Red circles correspond to inter-linker transitions in mixed ZIFs (there are two points for each mixed composition, corresponding to the two ordered configurations). Black circles refer to single-linker ZIFs.

the frontier orbitals of the linkers. We have compiled a larger list of modified imidazolate linkers, and calculated their HOMO and LUMO, which are provided in the Supporting Information. The motivation is that this list can serve as a reference for the electronic structures of functionalized ZIF linkers, and as a starting point for future design of mixed-linker ZIFs with tailored bands.

Finally, it is important to discuss how the electronic properties of the proposed ZIFs are modified by the presence of transition-metal cations (instead of Zn) in the tetrahedral sites. The established picture of photocatalysis by MOFs requires that photoexcitations involve ligand-to-metal charge transfer (LMCT),^[18] because this allows 1) strong light absorption due to the redox nature of the metal center, and 2) effective electron-hole separation to minimize recombination. While the mixed-linker Zn-based ZIFs proposed above exhibit a fundamental gap involving charge separation (from one kind of linker to the other), such excitation might not occur from optical absorption, because it would involve charge transfer through the Zn^{II} center, which does not have any empty levels at the same energy range as the LUCO. Furthermore, even if the inter-linker charge transfer in the Zn-based mixed ZIFs was possible, the absence of a metal redox center might lead to light absorption that is too weak for the system to work as an efficient photocatalyst.

We have therefore considered the cases of Co or Cu on the tetrahedral sites of the mIm/nIm mixed ZIF. These two metals are chosen because they are known from experiment to incorporate in ZIF structures, and in fact they constitute the metal centers in two reported ZIF photocatalysts.^[7] Figure 2d shows that the transition metals introduce empty d-levels above the LUCO level of the ZIF. In the case of Co, the empty d-band edge is too high, but in the case of Cu the empty d-band edge is only about 0.6 eV above the LUCO. In fact, in the Cu case the LUCO and the Cu 3d band clearly interact leading to some lowering of the LUCO energy. The presence of Cu and the overlap of its empty 3d levels with the LUCO of the ZIFs have important implications for photocatalysis. The photoadsorption leading to electron excitation would be stronger thanks to the $\text{Cu}^{\text{II}}/\text{Cu}^{\text{I}}$ redox pair. In addition, the

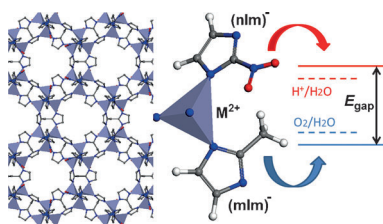
Communications



Zeolitic Imidazolate Frameworks

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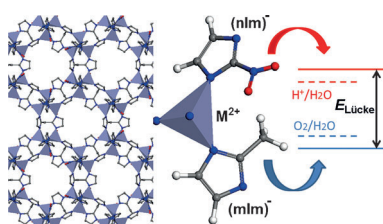
The sum of its parts: A computational study shows that using mixed linkers within zeolitic imidazolate frameworks provides a route for tuning their electronic gaps and band edge positions. It is theoretically possible in this way to match the electronic structure requirements for photocatalytic reactions including water splitting and CO₂ reduction.



Zeolithische Imidazolat-Gerüste

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Die Summe der Teile: Rechnungen zeigen, dass sich elektronische Lücken und die Lage der Bandkanten von zeolithischen Imidazolat-Gerüsten durch den Einsatz von Linkermischungen einstellen lassen. Theoretisch können die elektronischen Strukturen auf diese Weise für photokatalytische Reaktionen wie Wasserspaltung und CO₂-Reduktion abgestimmt werden.

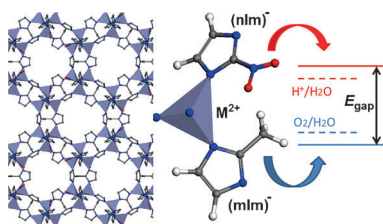
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